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## What is claimed is:

- 1. A method for identifying a polypeptide that binds a ligand, comprising:
- (a) comparing a sequence of a polypeptide 5 to a sequence model for polypeptides that bind a ligand, wherein said sequence model comprises representations of amino acids consisting of a subset of amino acids, said subset of amino acids having one or more atom within a selected distance from a bound ligand in said
- 10 polypeptides that bind said ligand; and
- (b) determining a relationship between said sequence and said sequence model, wherein a correspondence between said sequence and said sequence model identifies said polypeptide as a polypeptide that 15 binds said ligand.
  - 2. The method of claim 1, wherein said sequence model comprises a nucleic acid sequence.
  - 3. The method of claim 1, wherein said sequence model comprises an amino acid sequence.
- 4. The method of claim 1, wherein one of said sequence models is a Hidden Markov Model.
  - 5. The method of claim 1, wherein one of said sequence models is a Support Vector Machines Model.

- 6. The method of claim 1, wherein one of said sequence models is a Position Specific Score Matrices Model.
- 7. The method of claim 1, wherein one of said 5 sequence models is a Neural Network Model.
  - 8. The method of claim 1, further comprising the step of:
- (c) producing a sequence model with a set of sequences, said set of sequences consisting of 10 sequences of polypeptides having a subset of amino acids, said subset of amino acids having one or more atom within a selected distance from a bound ligand in said polypeptides that bind said ligand.
- 9. The method of claim 8, further comprising 15 the steps of:
  - (d) adding a sequence of said identified polypeptide that binds said ligand to said set of sequences; and
- (e) repeating steps (a) through (c) one or 20 more times.

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- 10. The method of claim 1, wherein said sequence model is produced by the steps of:
- (a) identifying a subset of amino acidshaving one or more atom within a selected distance from a5 bound conformation of a ligand in a set of polypeptidesthat bind said ligand; and
- (b) producing a sequence model, amino acids of said sequence model consisting of said subset of 10 amino acids.
  - 11. A method for identifying a member of a pharmacofamily, comprising:
- (a) comparing a sequence of a polypeptideto a sequence model for polypeptides of a pharmacofamily;and
- (b) determining a relationship between said sequence and said sequence model, wherein a correspondence between said sequence and said sequence model identifies said polypeptide as a member of said 20 pharmacofamily.
  - 12. The method of claim 11, wherein said sequence model comprises a nucleic acid sequence.
  - 13. The method of claim 11, wherein said sequence model comprises an amino acid sequence.
- 25 14. The method of claim 11, wherein said sequence model is a Hidden Markov Model.

- 15. The method of claim 11, wherein said sequence model is a Support Vector Machines Model.
- 16. The method of claim 11, wherein said sequence model is a Position Specific Score Matrices 5 Model.
  - 17. The method of claim 11, wherein one of said sequence models is a Neural Network Model.
  - 18. The method of claim 11, further comprising the step of:
- 10 (c) producing a sequence model with a set of sequences, said set of sequences consisting of sequences of polypeptides in said pharmacofamily.
  - 19. The method of claim 18, further comprising the steps of:
- (d) adding a sequence of said identified member of said pharmacofamily to said set of sequences; and
  - (e) repeating steps (a) through (c) one or more times.
- 20. The method of claim 11, wherein said sequence model comprises representations of amino acids consisting of a subset of amino acids, said subset of amino acids having one or more atom within a selected distance from a bound ligand in said polypeptides of said pharmacofamily.

- 21. The method of claim 20, wherein said sequence model is produced by the steps of:
- (a) identifying a subset of amino acids in a pharmacofamily having one or more atom within a5 selected distance from a bound conformation of a ligand; and
  - (b) producing a sequence model, amino acids of said sequence model consisting of said subset of amino acids.
- 22. A method for identifying a member of a pharmacofamily, comprising:
  - (a) comparing a sequence of a polypeptide to a sequence model and a differential sequence model; and
- 15 (b) determining a relationship between said sequence and said sequence models, wherein a correspondence between said sequence and said sequence models identifies said polypeptide as a member of said pharmacofamily.
- 20 23. The method of claim 22, wherein said sequence model comprises a nucleic acid sequence.
  - 24. The method of claim 22, wherein said sequence model comprises an amino acid sequence.
- 25. The method of claim 22, wherein one of 25 said sequence models is a Hidden Markov Model.

- 26. The method of claim 22, wherein one of said sequence models is a Support Vector Machines Model.
- 27. The method of claim 22, wherein one of said sequence models is a Position Specific Score
  5 Matrices Model.
  - 28. The method of claim 22, wherein one of said sequence models is a Neural Network Model.
  - 29. The method of claim 22, further comprising the step of:
- of sequences, said set of sequences consisting of sequences of polypeptides in said pharmacofamily.
  - 30. The method of claim 29, further comprising the steps of:
- (d) adding a sequence of said identified member of said pharmacofamily to said set of sequences; and
  - (e) repeating steps (a) through (c) one or more times.
- 31. The method of claim 22, wherein said differential sequence model comprises representations of amino acids consisting of a subset of amino acids, said subset of amino acids having one or more atom within a selected distance from a bound ligand in said
- 25 polypeptides of said pharmacofamily.



- 32. The method of claim 31, wherein said differential sequence model is produced by the steps of:
- (a) identifying a subset of amino acids in a pharmacofamily having one or more atom within a5 selected distance from a bound conformation of a ligand; and
  - (b) producing a differential sequence model, amino acids of said differential sequence model consisting of said subset of amino acids.

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